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Detailed Structural Investigation of the Grafting of [Ta(=CHtBu)(CH2tBu)3] and [Cp*TaMe4] on Silica Partially Dehydroxylated at 700 °C and the Activity of the Grafted **Complexes toward Alkane Metathesis**

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Abstract: The reaction of [Ta(=CHtBu)(CH2tBu)3] or [Cp*Ta(CH3)4] with a silica partially dehydroxylated at 700 °C gives the corresponding monosiloxy surface complexes [(≡SiO)Ta(=CHtBu)(CH₂tBu)₂] and [(≡SiO)Ta(CH₃)₃Cp*] by eliminating a σ-bonded ligand as the corresponding alkane (H−CH₂fBu or H−CH₃). EXAFS data show that an adjacent siloxane bridge of the surface plays the role of an extra surface ligand, which most likely stabilizes these complexes as in [(=SiO)Ta(=CHtBu)(CH₂tBu)₂(=SiOSi=)] (1a') and $[(\equiv SiO)Ta(CH_3)_3Cp^*(\equiv SiOSi\equiv)]$ (2a'). In the case of $[(\equiv SiO)Ta(\equiv CH_tBu)(CH_2tBu)_2(\equiv SiOSi\equiv)]$, the structure is further stabilized by an additional interaction: a C-H agostic bond as evidenced by the small J coupling constant for the carbenic C-H (J_{C-H} = 80 Hz), which was measured by J-resolved 2D solid-state NMR spectroscopy. The product selectivity in propane metathesis in the presence of [(=SiO)Ta(=CHtBu)-(CH₂tBu)₂(≡SiOSi≡)] (1a') as a catalyst precursor and the inactivity of the surface complex [(≡SiO)Ta-(CH₃)₃Cp*(≡SiOSi≡)] (2a') show that the active site is required to be highly electrophilic and probably involves a metallacyclobutane intermediate.

Introduction

The interaction of organometallic complexes with oxide surfaces such as silica or alumina has been studied for 30 years. 1-5 The first application of these materials was to the generation of highly active heterogeneous polymerization catalysts, followed by applications to other catalytic processes such as hydrogenation, olefin and alkane metathesis, Fischer Tropsch, or oxidation. This approach is referred to as surface organometallic chemistry (SOMC), and its main objective is the transfer of concepts and tools from molecular chemistry to surface science.⁶ The recent development of advanced spectro-

scopic techniques has greatly helped to characterize the exact 42 nature of active sites of these systems.^{7–10} Notably, we show how it is almost essential to combine the use of several analytical techniques to avoid making misleading conclusions. In this light, we discuss our results in comparison to previous studies carried out on these systems.¹¹ In the following study, we study and compare the reactivity of silica partially dehydroxylated at 700 $^{\circ}$ C with two tantalum complexes, [Ta(=CH $_t$ Bu)(CH $_2t$ Bu) $_3$] and [Cp*TaMe₄]. The structure of each surface complex and the role of the silica surface on these structures will be discussed, highlighting the advantage of a rigorous multistep approach to

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Table 1. Solid-State NMR Data for Solid 1

ligand	δ $^{1}\text{H/ppm}$	δ $^{13}{ m C/ppm}$	J_{C-H}/Hz
=СНСМе3	4.2	247	80
$=CHCMe_3$		47	
$=CHCMe_3$	1.0	31	126
$-\mathbf{CH}_2\mathbf{CMe}_3$	1.0	95	125
$-CH_2CMe_3$		31	
$-CH_2C\mathbf{Me}_3$	1.0	31	126

surface organometallic chemistry. The choice of tantalum for such a study has been based on its unusual reactivity toward alkanes because alkane metathesis, a reaction which transforms an alkane into its higher and lower homologues, was originally discovered with $[(\equiv SiO)_2Ta-H]$, a d² electron complex alkane metathesis catalyst.¹² We have recently disclosed that the mixture of $[(\equiv SiO)Ta(\equiv CHtBu)(CH_2tBu)_2]$ (1a) $[(\equiv SiO)_2 Ta(\equiv CHtBu)(CH_2tBu)]$ (1b), both d⁰ electron complexes, is also a catalyst precursor for this reaction.¹³ Because this study yielded two well-defined complexes, their activity in alkane metathesis was tested to gather information on the key requirements for an alkane metathesis catalyst precursors.

Results and Discussion

Reactivity of [Ta(=CHtBu)(CH2tBu)3] with Silica Partially Dehydroxylated at 700 °C. An exemplary case for multitechnique SOMC is given by the studies of the reactivity of [Ta(=CHtBu)(CH2tBu)3] with partially dehydroxylated silica.8,11,14,15 In the specific case of a silica partially dehydroxylated at 700 °C, the reaction leads to a yellow solid 1, whose structure can be formulated as the well-defined monosiloxy TaV carbene $[(\equiv SiO)Ta(\equiv CHtBu)(CH_2tBu)_2]$, 1a, as deduced by the combined use of mass balance analyses, IR, 1D and 2D HETCOR NMR spectroscopies (Table 1), and characterization through chemical reactivity studies (such as pseudo-Wittig or hydrolysis).

Furthermore, isotopic distribution studies¹⁴ and 1D solid-state NMR spectroscopy on selectively ¹³C-labeled complexes⁸ have indicated the involvement of the surface intermediate [(≡SiO)-Ta(CH₂tBu)₄], 1c, during the grafting reaction, and the slow transformation of 1c into $[(\equiv SiO)Ta(\equiv CHtBu)(CH_2tBu)_2]$ 1a and neopentane. The identification and characterization of 1c, and the related mechanism of the grafting reaction, was further substantiated by studies of the reaction of the organometallic precursor with a molecular model for silica's surface isolated grafting site, the polyhedral oligosilsesquioxane $[(c-C_5H_9)_7-$ Si₇O₁₂Si(OH)].⁸ Studies based solely on ¹³C solid-state NMR data for this reaction have postulated the same intermediate species, although NMR data have not always been correctly assigned.11

Ideally, one would like to obtain crystallographic data to yield the interatomic geometric parameters of the coordination sphere of a metal center and its surroundings in the same way molecular chemists use X-ray crystallography diffraction studies on single crystals. In the case of silica, an amorphous support, extended X-ray absorption fine structure (EXAFS) analysis provides

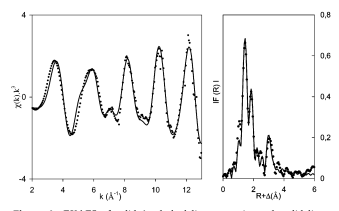


Figure 1. EXAFS of solid 1: dashed lines, experimental; solid lines, spherical wave theory.

Table 2. EXAFS Parameters for Solid 1a

neighboring atom	# of atoms ^b distance (/Å)		Debye-Waller factor (/Å)	
=CHCMe ₃	1	1.898(8)	0.07(6)	
− O Si	1	1.898	0.03(2)	
$-CH_2CMe_3$	2	2.150(4)	0.07(2)	
$-\mathbf{OSi}_2$	1	2.64(1)	0.11(4)	
-CH ₂ CMe ₃	3	3.417(5)	0.06(2)	

^a Fit residue: $\rho = 5.8\%$. ^b All shells fit with an overall scale factor of 1.1.

insight into the number and distances of first and second neighbors. The EXAFS data collected on the solid 1 are consistent with the following features (Figure 1 and Table 2): (i) two neighbors (either carbon or oxygen atoms) at a short distance (1.898 Å, not resolved), (ii) two other carbon neighbors at a longer distance (2.150 Å), (iii) an extra oxygen atom at a much longer distance (2.64 Å), and (iv) three carbon atoms at 3.417 Å. The two neighbors at a short bond distance (1.898 Å) can be assigned to an alkylidene (=CHtBu and a σ -bonded siloxy (**O**Si≡) substituent, while those at 2.150 Å are assigned to two σ -bonded carbons of the neopentyl groups (CH₂tBu). The extra O-atom neighbor at 2.64 Å is assigned to a siloxane bridge which acts as a two-electron donor ligand to stabilize the otherwise highly electron unsaturated surface complex 1a (formally a 10-electron complex) to yield the more stabilized 12-electron species [(≡SiO)Ta(=CHtBu)(CH2tBu)2- $(\equiv SiOSi \equiv)], 1a'.$

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The proposed ligand assignments for the observed bond distances are consistent with corresponding bond distances obtained by X-ray crystallography in analogous molecular complexes, such as $d(Ta=CHtBu) = 1.89 \text{ Å},^{16,17} d(Ta-C) =$ $2.19 \text{ Å},^{18} d(\text{Ta-OSi}) = 1.89 \text{ Å},^{19} \text{ and } d(\text{Ta-O}) = 2.25 - 2.35$ Å for a coordinated ether.^{20,21}

Recently, J-resolved 2D solid-state NMR spectroscopy was introduced as a novel method to measure the M-C-H bond angle and was applied to the study of surface-bound metallo-

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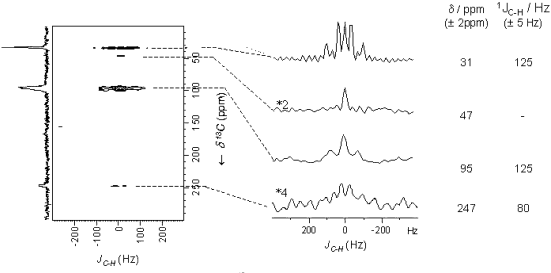


Figure 2. 2D J-resolved solid-state NMR spectrum of solid 1, 10% 13 C enriched at the α positions (*). Traces extracted along the ω_1 dimension of the 2D J-resolved spectrum at different carbon chemical shift frequencies: 31, 47, 95, and 247 ppm.

Scheme 1. Reactivity of [Ta(=CHfBu)(CH₂fBu)₃] with SiO₂-(700): Formation of [≡SiO-Ta(=CHfBu)(CH₂fBu)₂(≡SiOSi≡)] (1a')

carbenes⁹ because their J(C-H) coupling constant is strongly correlated with this angle. 16,22,23

The results of the J-resolved 2D solid-state NMR study of 1a' are reported in Table 2 and Figure 2.

The carbenic signal at 247 ppm appears as a doublet with a very small coupling constant of ${}^{1}J_{C-H} = 80$ Hz, strongly indicating that the C-H bond is stretched and that the Ta-C-H bond angle is very acute (probably lower than 90°). Similarly, the molecular silsesquioxane analogue, $[(c-C_5H_9)_7-$ Si₇O₁₂SiO)Ta(=CHtBu)(CH₂tBu)₂] (**1m**), also displays a doublet ($J_{C-H} = 86 \text{ Hz}$) at 245 ppm assigned to its carbonic carbons. Because similar spectroscopic features have been observed for the starting molecular complex, and explained in terms of an agostic interaction between the metallic center and the carbenic proton, 16,24-26 we propose that the same type of agostic interaction is also present for the surface-bound Ta center and its α -carbenic proton of 1a'.

The signal at 95 ppm, assigned to methylene (CH₂tBu), is a triplet, with C-H coupling constant ${}^{1}J_{C-H} = 125$ Hz, as expected for an sp³ carbon. The molecular complex 1m exhibits the analogous signal at 98.7 ppm (triplet, ${}^{1}J_{C-H} = 109$ Hz). In addition, there is a sharp singlet at 40 ppm, which can be assigned to the quaternary carbon of the tertiobutyl group attached to the carbene ligand; one other multiplet can be observed at 30 ppm, this resonance corresponding probably to the superimposition of a quadruplet (${}^{1}J_{C-H} = 125 \text{ Hz}$) due to the methyl signals and a singlet due to the quaternary carbon of the neopentyl ligands.

The combined use of EXAFS, IR, mass balance analyses, NMR studies (including ${}^{1}J_{C-H}$ -resolved data), and studies with molecular analogues yields a sharp description of the grafting reaction of organometallic precursor [Ta(=CHtBu)(CH2tBu)3] on a silica surface (see Scheme 1).

Particularly noteworthy, the EXAFS and J-resolved studies have highlighted the presence of stabilizing interactions at the Ta-center, beside the σ -bonded alkyl and alkylidene ligands, with which the tantalum center (formally a tetracoordinated 10electron species, 1a) achieves a pseudo-octahedral geometry through (i) a 2-electron donor interaction with an adjacent (≡SiOSi≡) surface bridge, and (ii) an agostic interaction with the carbenic C-H bond, to yield the stabilized formally 14electron species, $[(\equiv SiO)Ta(\equiv CHtBu)(CH_2tBu)_2(\equiv SiOSi\equiv)]$, 1a'.

Reactivity of [Cp*TaMe4] with Silica Partially Dehydroxylated at 700 °C. The reports available on the reactivity of Cp*TaMe4 (Scheme 2) with partially dehydroxylated silica are comprised of spectroscopic ¹H- and ¹³C-CP NMR studies. ¹¹ The data suggest that the organometallic surface species [(≡SiO)TaCp*Me₃], 2a, is obtained, either by reaction with surface silanol [≡SiOH] and elimination of methane, or by cleavage of a surface siloxy bridge [≡SiOSi≡] and formation 148

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Scheme 2. Reactivity of $[Cp^*Ta(CH_3)_4]$ with $SiO_{2-(700)}$: Formation of $[\equiv SiO-Ta(CH_3)_3Cp^*(\equiv SiOSi\equiv)]$ (2a')

of silicon bound methyl moiety, [≡Si-Me]. The presence of this latter species is inferred by one signal at −6 ppm in the ¹³C NMR spectrum.¹¹

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Herein is discussed in detail the reaction of [Cp*TaMe₄] with a silica partially dehydroxylated at 700 °C. First, a silica disk partially dehydroxylated at 700 °C (SiO₂₋₍₇₀₀₎) was immersed at room temperature in a yellow pentane solution of [Cp*TaMe₄]-(1.2 equiv/surface silanols; 0.26 \pm 0.03 mmol accessible OH/g). After the excess molecular complex was washed and the resulting solid was dried under dynamic vacuum (10⁻⁵ Torr, 2 h), an IR spectrum was recorded (Figure 3).

The narrow band assigned to isolated silanols at 3747 cm⁻¹ totally disappeared, leaving a broad band in the 3740–3550 cm⁻¹ region as well as two sets of bands in the 3000–2700 and 1500–1300 cm⁻¹ regions. The broad band corresponds to $\nu_{\rm (O-H)}$ of residual silanols in interaction with the perhydrocarbyl ligands present at the surface of silica (vide infra for further comments), the two latter sets of IR bands being assigned to $\nu_{\rm (CH)}$ and $\nu_{\rm (C=C)}/\delta_{\rm (CH)}$ vibrations.

Second, using larger quantities of silica (0.20-1.0 g) allowed the methane released to be quantified: 1.1-1.2 equiv of methane was evolved per grafted Ta, consistent with a chemical grafting of the molecular complex. This reaction occurs via cleavage of one (Ta-CH₃) bond by a surface silanol forming a covalent (Ta-Osi≡) bond and methane to yield a yellow solid 2. The formation of [(≡SiO)TaCp*Me₃], 2a, leads to 1 mol of MeH/ mol of grafted Ta as expected, and therefore small quantities of the bisiloxy species [(≡SiO)₂TaCp*Me₂], **2b**, for which 2 mol of MeH/mol of grafted Ta are expected, could also be possibly formed. Elemental analysis of the yellow solid shows the presence of 12 ± 2 carbons/Ta, which is consistent with the proposed structure (13 C/Ta and 12 C/Ta for 2a and 2b, respectively). The tantalum loading typically varies between 2.6 and 3.7 Ta % wt, depending on the batch of SiO₂₋₍₇₀₀₎ used (depending on how much silica was compacted). Such tantalum loadings correspond to 0.14-0.21 mmol of Ta/g of solid, while $SiO_{2-(700)}$ typically contains 0.26 \pm 0.03 mmol of OH/g on silica. Therefore, such low loading indicates a partial consumption of the surface silanols as already evidenced by in situ IR experiments (vide supra) where a broad band assigned to residual silanols was detected. This is in contrast to what has been observed in the reaction of [Ta(=CHtBu)(CH2tBu)3] with SiO₂₋₍₇₀₀₎, ¹⁵ for which all surface silanols are consumed, but it is consistent with the larger size of the [TaCp*Me₃] fragment as compared to that of [Ta(=CHtBu)(CH2tBu)2], which, in turn, would prevent the access of the molecular complex, [Cp*TaMe₄], to the remaining surface silanols.

Third, the solid-state ¹H MAS NMR spectrum of the yellow solid **2** shows an intense peak at 1.61 ppm along with broad signals between -0.3 and 0.7 ppm, as does the ¹H spectrum of

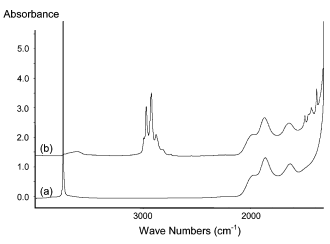


Figure 3. IR spectra of the reaction of $[Cp*Ta(CH_3)_4]$ with $SiO_{2-(700)}$ by IR spectroscopy: (a) $(SiO_{2-(700)})$ and (b) $[\equiv SiO-Ta(CH_3)_3Cp*-(\equiv SiOSi\equiv)]$.

its ¹³C-labeled (20% on the Me substituents attached to Ta) analogue, 2* (Figure 4a). The ¹³C CP/MAS NMR spectrum of 2 displays three signals at 117, 58, and 9 ppm, as does the spectrum of 2* (albeit with different relative ratios, Figure 4b). In accord with the literature data, 11 the signals at 117 and 9 ppm can be tentatively assigned to the carbons of the cyclopentadienyl ring and the methyl groups of the Cp* ligand, respectively, and the signal at 58 ppm can be assigned to that of the (Ta-Me₃) groups, which is shifted upfield as compared to that of the starting molecular complex [Cp*TaMe₄] (Ta-Me₄, 74 ppm). The large upfield shift in going from a molecular to a supported complex is consistent with the replacement of one methyl group by an electronegative siloxy group.^{27,28} No resonance of any significance could be detected at -6 ppm, which is the signature peak of [≡Si-Me] as previously reported,¹¹ and no further evidence for a bisiloxy species could be obtained through these NMR studies.

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Fourth, the 2D ¹H-¹³C HECTOR NMR spectrum of 2*, recorded with a contact time of 1 ms (Figure 4c), shows a correlation between the signal at 1.6 ppm in the F_1 dimension (1 H) and the signal at 9 ppm in the F_2 dimension (13 C), which are assigned to the methyl groups of the pentamethylcyclopentadienyl ring. The ¹³C resonance of the methyl groups directly bonded to the tantalum gives two pairs of correlations: -0.05ppm/58.0 ppm and -0.28 ppm/57.5 ppm in the F_1/F_2 dimensions. These two types of proton—carbon resonances probably correspond to the methyl group trans to oxygen (58.0 and -0.05ppm), and the other correlation (57.5 and -0.28 ppm) corresponds to the cis methyl groups as observed for similar molecular complexes, $Cp*TaXMe_3$ (X = Cl, OMe, OiPr, OtBu, NMe₂), which adopt a "four-legged piano stool" geometry and in which the two types of methyl groups are not equivalent.²⁷ The signal at 0.6 ppm in the F_1 dimension (1 H) does not correlate with any carbon (in the F_2 dimension), and this signal disappears when using deuterated silica (see Supporting Information). Therefore, the resonance at 0.6 ppm is most likely due to a surface silanol [≡SiO-H] resonance, which is shifted upfield

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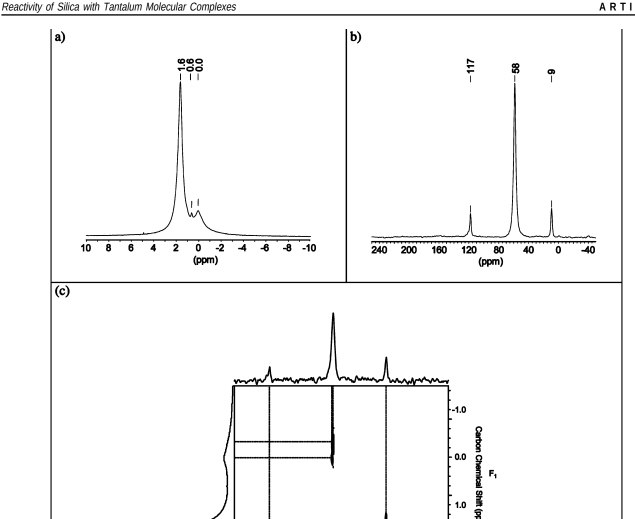


Figure 4. (a) ¹H MAS NMR spectrum of 2* (number of scans = 8, repetition delay = 8 s, 90° H pulse = 3 μs, no apodization). (b) ¹³C CP/MAS spectrum of 2* (number of scans = 78 339, repetition delay = 2 s, P15 = 5 ms, line broadening = 100 Hz). (c) 2D HETCOR solid-state NMR spectroscopy on 2*. The displayed spectra correspond to a ¹H MAS NMR spectrum (number of scans = 16, repetition delay = 8 s, line broadening = 5 Hz) for F₁ and ¹³C CP/MAS spectrum (number of scans = 1024, repetition delay = 2 s, P15 = 1 ms, line broadening = 80 Hz) for F_2 .

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Carbon Chemical Shift (ppm) F₂

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with respect to the silanols of silica (typically observed at 1.8 ppm), most likely because of a ring-current effect of the adjacent Cp* ring.

Finally, Tantalum L_{III}-edge EXAFS measurements on 2 provided further insight into the structure of this silica-supported Ta species (Figure 5).

The data are consistent with an average of 1.3 oxygen atoms at 1.931 Å, 2.7 carbon atoms at 2.142 Å, and 5.0 carbon atoms at 2.456 Å in the coordination sphere of Ta, which is in turn consistent with 2a being the major species present in the solid, and **2b** (if any) as a minor product (Table 3).

The proposed assignment for measured Ta-C bond distances is in good agreement with values obtained from crystallographic data for Ta-C(Me) (2.074-2.150 Å in [TaMe₅];²⁹ 2.115 Å in [TaMe₃(OAr)₂];³⁰ 2.181 Å in [Cp*TaMe₂(C₆H₄)];³¹ and 2.22-2.26 Å in $[(TaMe_3Cp^*)_2(\mu-O)]^{32})$ and Ta-C(pentamethylcyclopentadienyl) distances (2.345–2.406 Å in [Cp*TaCl₂Me₂];²⁸ 2.424-2.480 Å in [Cp*Ta(*p-tert*-butylcalix[4]arene)];³³ 2.366-2.518 Å in $[Cp*Ta(=NAr)Cl_2]^{34}$). Moreover, the agreement with the EXAFS data was improved when the model included five carbons at 3.44 Å, corresponding to the five methyl groups of the Cp* ligand $(3.42-3.61 \text{ Å in } [\text{Cp*Ta}(=\text{NAr})\text{Cl}_2]^{34})$, and an extra 1.3 oxygen atoms at 3.04 Å, which can be assigned to that of siloxane bridges close to a tantalum species, to yield

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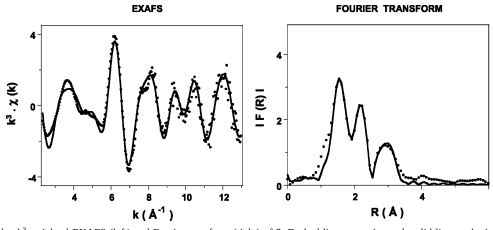


Figure 5. Ta L_{III}-edge k³-weighted EXAFS (left) and Fourier transform (right) of 2. Dashed lines, experimental; solid lines, spherical wave theory.

Table 3. EXAFS Parameters for Solid 2a

neighboring atom	# of atoms	distance from Ta (/Å)	Debye-Waller factor (/Å)
OSi	1.3	1.931	0.042
CH_3	2.7	2.142	0.097
C' (Cp* ring)	5.0	2.456	0.086
OSi ₂	1.3	3.040	0.063
C" (Me of Cp*)	4.8	3.445	0.082

^a Fit residue: $\rho = 7.8\%$.

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[(\equiv SiO)TaCp*Me₃(\equiv SiOSi \equiv)], **2a**′, analogous to the observations for the carbenic derivative discussed above [(\equiv SiO)Ta-(\equiv CH $_2$ tBu)(CH $_2$ tBu)2(\equiv SiOSi \equiv)], **1a**′.

In summary, the combined analysis of the experimental data collected on the reaction of Cp^*TaMe_4 with $SiO_{2-(700)}$ indicates the presence of $[(\equiv SiO)TaMe_3Cp^*(\equiv SiOSi\equiv)]$ (2a') as a major species, in which a siloxane bridge acts as an extra donor ligand to the Ta center. In contrast to most organometallic complexes studied so far, only part of the surface silanols reacted with $[Cp^*TaMe_4]$, as demonstrated by elemental analysis, IR, and NMR spectroscopies. The remaining silanols interact with a nearby Cp^* as observed through the upfield shift of $\delta(\equiv SiOH)$ in the 1H NMR spectrum. The grafting mechanism corresponds to the silanolysis of a Ta—Me bond by a surface $[\equiv SiOH]$, with elimination of 1 mole of methane, while no evidence was obtained supporting the previously proposed addition of a Ta—C across a Si—O bond of a surface siloxy bridge $[\equiv SiOSi\equiv]$. 11

Comparison of the Reactivity of [Cp*TaMe4] and [Ta(=CHtBu)(CH2tBu)3] Supported on Silica Partially Dehydroxylated at 700 °C toward Propane. In the case of $[(\equiv SiO)Ta(CH_3)_3Cp*(\equiv SiOSi\equiv)]$ (2a'), a sterically crowded electron-rich TaV surface complex, its reaction with propane yields methane as the sole gaseous product, which probably formed via the decomposition of 2a' (Table 4). On the other hand, $[(\equiv SiO)Ta(\equiv CHtBu)(CH_2tBu)_2(\equiv SiOSi\equiv)]$ (1a'), an electron-deficient TaV surface complex as evidenced by the C-H agostic interaction, catalytically transforms propane into its higher and lower homologues (Table 4). Moreover, initiation products, which result from the reaction of the neopentyl/ neopentylidene fragments in 1a' and propane, are also formed: 2,2-dimethylpropane (tBuCH₂-H, 1.05 equiv/Ta), 2,2-dimethylbutane (tBuCH₂-CH₃, 0.30 equiv/Ta), and 2,2-dimethylpentane (tBuCH₂-CH₂CH₃, 0.11 equiv/Ta). Note that no tBuCH₂- $CH_2CH_2CH_3$ (<0.1%, not detected) is formed. The ratio of initiation products tBuCH₂-CH₃/tBuCH₂-CH₂CH₃ is 2.5. Simi-

Table 4. Reactivity of Tantalum Species Supported on Silica toward Propane

catalysts	% wt	P	ratio	conv. ^a	TON ^a
	Ta	(Torr)	(nC ₃ H ₈ /nTa)	(%)	(mol P/mol Ta)
solid 2	3.5	495	805	0	$0 \\ 33-34^b \\ 65^c$
solid 1	3.9	600	580	5.8	
[(≡SiO) ₂ Ta−H]	5.0	550	1065	6.1	

 a As measured after 120 h. b The selectivities in methane, ethane, isobutane, butane, isopentane, pentane, and hexanes are 12.8%, 47.7%, 10.2%, 22.2%, 2.5%, 3.6%, and 0.9%, respectively. c The selectivities in methane, ethane, isobutane, butane, isopentane, pentane, and hexanes are 18.0%, 40.0%, 8.6%, 24.4%, 2.9%, 5.1%, and 1%, respectively.

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larly in olefin metathesis (Scheme 3), we have observed the initiation products of the reaction of propene and [(≡SiO)Re-(≡CtBu)(≔CHtBu)(CH2tBu)(≡SiOSi≡)]: tBuCH=CH2 and trans-tBuCH=CHCH3 (no cis tBuCH=CHCH3 is detected). 35,36 Their ratio (tBuCH=CH2/trans-tBuCH=CHCH3) is 3.0, which is closely related to that observed for the initiation products in the metathesis of propane on [(≡SiO)Ta(=CHtBu)(CH2tBu)2-(≡SiOSi≡)] (1a′). In this latter case, the selectivity in initiation products can be understood in terms of minimization of the steric interactions in the metallacyclobutane intermediates (or in their formation), 37 which is governed by the relative position of the substituents: typically substituents in the [1,3]-positions (usually both in equatorial positions) of the metallacyclobutane intermediate generate less steric hindrance than those in the [1,2]-positions (usually both in equatorial positions, Scheme 3). 38

The selectivity in propane metathesis can also be explained by using the same model in which [1,3]- and [1,2]-interactions determine the ratio of products. For instance, the butane/pentane ratios are 6.2 and 4.8 for **1a**′ and [(≡SiO)₂Ta−H], respectively (Table 4).³⁹ A similar trend is observed for the isobutane/ isopentane ratios, which are 4.1 and 3.0, respectively. The higher selectivity in butanes (the transfer of one carbon via metallacyclobutanes involving [1,3]-interactions) than that of pentanes

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Scheme 3. Possible Relation between the Mechanisms of Olefin and Alkane Metatheses: The Initiation Step

Scheme 4. Model for Product Selectivities in Alkane Metathesis

(the transfer of two carbons via metallacyclobutanes involving [1,2]-interactions) is consistent with this model (Schemes 3 and 4).

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Because alkane metathesis catalysts need to be highly electrophilic and coordinatively unsatured (Cp* is detrimental to catalysis), because the selectivities of initiation products and alkane metathesis products are similar to that observed for olefin metathesis, and because 1a', a metallocarbene, is a catalyst precursor for alkane metathesis, we therefore propose that one of the key steps in alkane metathesis would be the formation of a metallacyclobutane intermediate and probably involves

metallocarbenes rather than direct C-C σ-bond metathesis or oxidative addition pathways as was suggested earlier as a possible working hypothesis.

Experimental Details

General Procedure. All experiments were carried out under a controlled atmosphere, using Schlenk and glovebox techniques for the organometallic synthesis. For the synthesis and treatments of the surface species, reactions were carried out using high-vacuum lines (1.34 Pa) and glovebox techniques. SiO₂ (Aerosil Degussa, 200 m² g⁻¹) was compacted with distilled water, calcined (500 °C under air for 4 h),

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and partially dehydroxylated under vacuum (1.34 Pa) at 500 °C for 12 h and then at 700 °C for 4 h (support referred to as SiO₂₋₍₇₀₀₎). [Ta-(=CHtBu)(CH₂tBu)₃] and [Cp*Ta(CH₃)₃] were prepared according to the literature procedure. ⁴⁰ The 10% ¹³C-labeled [Ta(=CHtBu)-(CH₂tBu)₃] was prepared as reported previously. ^{8,10} ¹³C-labeled Cp*Ta-(CH₃)₄ was prepared following the literature procedure for the unlabeled complex, using ¹³CH₃Li prepared from Li wire and CH₃I. ⁴¹ Pentane and diethyl ether were distilled on NaK alloy and Na/benzophenone, respectively followed by degassing through freeze—pump—thaw cycles. Infrared spectra were recorded on a Nicolet Magna 550 FT spectrometer equipped with a cell designed for in situ reactions under controlled atmosphere. Elemental analyses were performed at the Service Central d'Analyses of CNRS in Solaize.

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¹H MAS and ¹³C CP-MAS solid-state NMR spectra were recorded on a Bruker DSX-300 spectrometer. For specific studies (see below), ¹H MAS and ¹³C CP-MAS solid-state NMR spectra were recorded on Bruker Avance-500 spectrometers with a conventional double resonance 4 mm CP-MAS probe at the Laboratoire de Chimie in Ecole Normale Supérieure de Lyon or at the Laboratoire de Chimie Organometallique de Surface in Ecole Supérieure de Chimie Physique Electronique de Lyon. The samples were introduced under Ar in a zirconia rotor, which was then tightly closed. In all experiments, the rotation frequency was set to 10 kHz unless otherwise specified. Chemical shifts were given with respect to TMS as external references for ¹H and ¹³C NMR.

Heteronuclear Correlation Spectroscopy. The two-dimensional heteronuclear correlation experiment was performed according to the following scheme: 90° proton pulse, t_1 evolution period, crosspolarization (CP) to carbon spins, detection of carbon magnetization. For the CP step, a ramp radio frequency (RF) field^{42,43} centered at 60 kHz was applied on protons, while the carbon RF field was matched to obtain optimal signal. The contact time for CP was set to 1 ms. During acquisition, the proton decoupling field strength was set to 83 kHz (TPPM decoupling⁴⁴). A total of 32 t_1 increments with 1024 scans each were collected. The spinning frequency was 10 kHz, and the recycle delay was 1 s (total acquisition time of 9 h). Quadrature detection in $ω_1$ was achieved using the TPPI method.⁴⁵

J-Resolved Spectroscopy. The two-dimensional J-resolved experiment was performed as previously described:9 after cross-polarization from protons, carbon magnetization evolves during t_1 under proton homonuclear decoupling. Simultaneous 180° carbon and proton pulses are applied in the middle of t_1 to refocus the carbon chemical shift evolution while retaining the modulation by the heteronuclear J_{CH} scalar couplings. A Z-filter is finally applied to allow phase-sensitive detection in ω_1 . Proton homonuclear decoupling was performed by using the frequency-switched Lee-Goldburg (FSLG) decoupling sequence. 46,47 Quadrature detection in ω_1 was achieved using the TPPI method.⁴⁵ The rotor spinning frequency was 10.2 kHz to synchronize the t1 increment with the rotor period. The proton RF field strength was set to 83 kHz during t₁ (FSLG decoupling) and acquisition (TPPM decoupling).⁴⁴ The lengths of carbon and proton 180° pulses were 7 and 6 μ s, respectively. An experimental scaling factor, measured as already described, 48 of 0.52 was found, which gave a corrected spectral width of 2452 Hz in 417

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Extended X-ray Absorption Fine Structure Spectroscopy (**EXAFS**). EXAFS on **1a** was carried out at the Stanford Synchrotron Radiation Laboratory (SSRL) at beamline 4-1 as previously reported. 10 Samples 2 was packaged within an argon-filled drybox in double airtight sample holders equipped with Kapton windows. X-ray absorption spectra were acquired at the Laboratoire pour l'Utilisation du Rayonnement Electromagnétique (LURE in Orsay, France), on the DCI ring at beam line D44. They were recorded at room temperature at the tantalum L_{III} edge, from 9700 to 11 000 eV, with a 2 eV step in the transmission mode. The data analysis was performed by standard procedures using either the suite of programs EXAFSPAK developed by G. George of SSRL or the one developed by A. Michalowicz.⁴⁹ Each spectrum was carefully extracted, and the best removal of lowfrequency noise was checked by further Fourier transformation. Fitting of the spectrum was done on the k^3 weighted data using the following EXAFS equation where S_0^2 is the scale factor; N_i is the coordination number of shell i; S_i is the central atom loss factor for atom i; F_i is the EXAFS scattering function for atom i; R_i is the distance to atom i from the absorbing atom; λ_i is the photoelectron mean free path; σ_i is the Debye-Waller factor; ϕ_i is the EXAFS phase function for atom i; and ϕ_c is the EXAFS phase function for the absorbing atom.

$$\chi(k) \simeq S_0^2 \sum_{i=1}^n \frac{N_i S_i(k, R_i) F_i(k, R_i)}{k R_i^2} \exp\left(\frac{-2R_i}{\lambda(k, R_i)}\right) \times \exp(-2\sigma_i^2 k^2) \sin[2kR_i + \phi_i(k, R_i) + \phi_c(k)]$$

The program FEFF7⁵⁰ was used to calculate theoretical values for S_i , F_i , λ_i , ϕ_i , and ϕ_c based on model clusters of atoms in which atomic positions were taken from the crystal structure of the most similar complexes. The refinements were performed by fitting the structural parameters N_i , R_i and σ_i . The fit residue, ρ , was calculated by the following formula:

$$\rho = \frac{\sum_{k} [k^{3} \chi_{\text{exp}}(k) - k^{3} \chi_{\text{Cal}}(k)]^{2}}{\sum_{k} [k^{3} \chi_{\text{exp}}(k)]^{2}}$$

Reaction of Silica Partially Dehydroxylated at 700 °C with [Ta-(CHtBu)(CH2tBu)₃], Formation of the Solid 1. A mixture of [Ta-(CHtBu)(CH2tBu)₃] (0.155 g, 0.33 mmol) in pentane (10 mL) and $SiO_{2-(700)}$ (1.0 g) was stirred at 25 °C for 2 h. After filtration, the solid was washed three times with pentane. The solvent was then removed, and the yellow orange solid was dried under dynamic-vacuum at 25 °C.

Reaction of Silica Partially Dehydroxylated at 700 °C with [Cp*TaMe₄]: Formation of the Solid 2. A mixture of Cp*TaMe₄ (42.5 mg, 0.11 mmol, 1.2 equiv) and $SiO_{2-(700)}$ (373 mg) in pentane (5 mL) was stirred at 25 °C for 2 h. After filtration, the solid was washed three times with pentane and all volatile compounds were condensed into another reactor (of known volume) to quantify methane evolved during the grafting. The resulting yellow powder was dried under vacuum (1.34 Pa) to yield 406 mg of 2. Analysis by gas chromatrography indicated the formation of 86 μ mol of methane during the grafting (1.0 μ CH₄/ μ Ta).

Reactivity of Propane on Solid 2. In a 352 mL volume reactor were added 67 mg of solid **2** (12.9 μ mol of Ta, 3.5%_{Ta}) and propane

the ω_1 dimension. The recycle delay was 1.3 s, and a total of 80 t_1 increments with 1024 scans each were collected (total acquisition time = 30 h).

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Reactivity of Silica with Tantalum Molecular Complexes

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(495 Torr, 9.38 mmol). The reaction mixture was heated at 150 °C for 120 h, during which small aliquots were analyzed by GC and GC/MS.

Reactivity of Propane on Solid 1. In a 235 mL volume reactor were added 60 mg of solid 1 (12.9 μ mol Ta, 3.9%_{Ta}) and propane (600 Torr, 7.59 mmol). The reaction mixture was heated at 150 °C for 120 h, during which small aliquots were analyzed by GC and GC/MS.

Conclusion

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Generally, whether using [Ta(=CHtBu)(CH2tBu)3] or [Cp*Ta-(CH₃)₄], the reaction with a silica partially dehydroxylated at 700 °C provides primarily the corresponding monosiloxy surface complexes $[(\equiv SiO)Ta(\equiv CHtBu)(CH_2tBu)_2(\equiv SiOSi\equiv)]$ (1a') and $[(\equiv SiO)Ta(CH_3)_3Cp^*(\equiv SiOSi\equiv)]$ (2a'), by eliminating a σ-bonded ligand as the corresponding alkane (H-CH₂tBu or H-CH₃). Moreover, when the metal is grafted, the coordination number is increased by coordination of a pair of electrons from the siloxane bridge, probably to stabilize the structure. In particular, in the case of $[Ta(=CHtBu)(CH_2tBu)_3]$, the surface silanol [≡SiOH] reacts preferentially with the carbene, to yield $[(\equiv SiO)Ta(\equiv CHtBu)(CH_2tBu)_2(\equiv SiOSi\equiv)]$, which is further stabilized by an additional C-H agostic interaction. In the case of [Cp*Ta(CH₃)₄], the size of the complex is such that after grafting some surface silanols do not become accessible but interact with adjacent Cp* ligands as evidenced by a strong upfield shift of this signal.

In conclusion, the combined evidence presented here shows the possibility for silica partially dehydroxylated at 700 °C to act as a LX ligand in the Green formalism, 3,51,52 rather than an X ligand, as is the most commonly reported. Moreover, the formation of well-defined species characterized at a molecular

level provides the possibility for one to probe mechanisms in heterogeneous catalysis through structure—activity relationship. In particular, in the case of alkane metathesis, the absence of activity of $[(\equiv SiO)Ta(CH_3)_3Cp*(\equiv SiOSi\equiv)]$ (2a') and the product selectivity when $[(\equiv SiO)Ta(\equiv CHtBu)(CH_2tBu)_2$ (≡SiOSi≡)] (1a') was used as a catalyst precursor show that the active site is required to be highly electrophilic, coordinatively unsaturated, and probably involves a metallacyclobutane intermediate. We are currently trying to probe the other steps of the mechanisms such as how the alkane is activated and how the olefins are formed.

DATE: August 26, 2004

This insight was gained by the combined use of several analytical techniques, with particular relevance given to the necessity to complement more traditional spectroscopic data (IR, one-dimensional solid-state NMR...) with data from techniques such as EXAFS and multidimensional NMR spectroscopies.

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Supporting Information Available: NMR spectra. This material is available free of charge via the Internet at http://pubs.acs.org.

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